## WHAT IS CLAIMED IS:

1. A pharmaceutical composition comprising a molecule comprising a fucose group in an  $\alpha$ 1,2 linkage, an  $\alpha$ 1,3 linkage or an  $\alpha$ 1,4 linkage to a galactose group and a pharmaceutically acceptable carrier.

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2. The composition of claim 1 where in the fucose is contained within an LNF-I group, an 2'FL group, an LNF-I group, an LNF-II group, an 3'FL group, an LNF-III group, an LDFH-I group, a LDFT group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.

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- 3. The composition of any of the forgoing claims wherein the molecule is a glycan, a glycolipid, a glycoprotein, a glycosaminoglycan or a mucin.
- 4. The composition of any of the forgoing claims wherein the molecule

  comprises at least two different groups selected from an LNF-I group, an 2'FL group, an

  LNF-I group, an LNF-II group, an 3'FL group, an LNF-III group, an LDFH-I group, a LDFT

  group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.
  - 5. The composition of any of the forgoing claims wherein the molecule comprises at least three different groups selected from an LNF-I group, an 2'FL group, an LNF-II group, an LNF-II group, an 3'FL group, an LNF-III group, an LDFH-I group, a LDFT group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc. .
- 6. The composition of any of the forgoing claims wherein the molecule contains at least 5 groups selected from an LNF-I group, an 2'FL group, an LNF-I group, an LNF-II group, an 3'FL group, an LNF-III group, an LDFH-I group, a LDFT group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.
- 7. The composition of any of the forgoing claims wherein the groups are covalently linked to a protein in an O-link to Ser or Thr or an N-link to Asn.

8. The composition of any of the forgoing claims wherein the composition does not contain a mammalian milk.

- 9. The composition of any of the forgoing claims wherein the composition does not contain human milk
  - 10. A pharmaceutical composition comprising a purified protein modified to include at least two different groups selected from:

10 2'-Fucosyllactose;

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Lacto-N-fucopentaose I;

Lacto-N-fucopentaose II;

3'-Fucosyllactose;

Lacto-N-fucopentaose II;

Lacto-N-difucohexaose I;

Lactodifucotetraose;

LactoN-tetraose;

LactoN-neotetraose;

3'-Sialyllactose;

3'-Sialyllactosamine;

6'-Sialyllactose;

6'-Sialyllactosamine;

Sialyllacto-N-neotetraose c;

Monosialyllacto-N-hexaose;

Disialyllacto-N-hexaose I;

Monosialyllacto-N-neohexaose I;

Monosialyllacto-N-neohexaose II

Disialyllacto-N-neohexaose

Disialyllacto-N-tetraose;

Disialyllacto -N-hexaose II;

Sialyllacto-N-tetraose a;

Disialyllacto-N-hexaose I;

Sialyllacto-N-tetraose b;

3'-Sialyl-3-fucosyllactose;

Disialomonofucosyllacto-N-neohexaose;

Monofucosylmonosialyllacto-N-octaose (sialyl Lea);

Sialyllacto-N-fucohexaose II;

Disialyllacto-N-fucopentaose II;

Monofucosyldisialyllacto-N-tetraose, or a variant thereof wherein Glc at the reducing end is replaced with GlcNAc.

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11. A pharmaceutical composition comprising a purified protein modified to include at least two different groups selected from:

2-Fucosyllactose;

Lacto-N-fucopentaose I;

Lacto-N-fucopentaose II;

3-Fucosyllactose;

Lacto-N-fucopentaose II;

Lacto-N-difucohexaose I;

Lactodifucotetraose; or a variant thereof wherein Glc at the reducing end is replaced with GlcNAc.

- 12. The composition of any of claims 10-11 wherein the protein is modified to contain multiple copies of each of the at least two different groups.
- 13. The composition of any of claims 10-12 wherein the protein is a human milk protein.
- 14. The composition of claims 10-13 wherein the human milk protein is selected from: κ-casein, α-lactalbumin, lactoferrin, bile salt-stimulated lipase, lysozyme, serum

albumin, folate-binding protein, haptocorrin, lipoprotein lipase, glycosaminoglycan, mucin, lactoperoxidase, and amylase.

- 15. The composition of any of the forgoing claims which is a synthetic composition.
  - 16. The composition of any of the forgoing claims which is not mammalian milk.
- 17. The composition of any of the forgoing claims further comprising at least one vitamin.
  - 118. The composition of any of the forgoing claims further comprising at least one mineral.
- 19. The composition of of any of the forgoing claims further comprising at least one edible fat.
  - 20. A pharmaceutical composition comprising a purified protein modified to include at least two different groups selected from:

20 2'-Fucosyllactose;

Lacto-N-fucopentaose I;

Lacto-N-fucopentaose II;

3'-Fucosyllactose;

Lacto-N-fucopentaose II;

Lacto-N-difucohexaose I;

Lactodifucotetraose;

2'-FLNac, or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc;

wherein the protein is not modified to contain any other oligosaccarides.

21. A synthetic nutritional composition comprising a glycan, a glycolipid, a glycoprotein, a glycosaminoglycan or a mucin that comprises at least two different groups selected from an LNF-I group, and 2'FL group, an LDFH-I group and a LDFT group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.

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22. The synthetic nutritional composition of claim 21 wherein the molecule comprises at least three different groups selected from an LNF-I group, and 2'FL group, an LDFH-I group and a LDFT group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.

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- 23. The synthetic nutritional composition of any of claims 21-22 wherein the molecule contains multiple copies of each of the at least two different groups.
- 24. The synthetic nutritional composition of any claims 21-23 wherein the molecule contains at least two copies of each of the at least two different groups.
  - 25. The synthetic nutritional composition of any of claims 21-24 wherein the molecule contains at least five copies of each of the at least two different groups.
- 26. The synthetic nutritional composition of any claims 21-25 wherein the molecule contains at least ten copies of each of the at least two different groups.
  - 27. The synthetic nutritional composition of any claims 21-26 wherein the molecule contains at least 20 copies of each of the at least two different groups.

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28. A synthetic nutrition composition comprising a purified protein modified to include a group selected from: a Lacto-N-fucopentaose I group, a Lacto-N-fucopentaose II group, a 2-Fucosyllactose group, a 3-Fucosyllactose group, a Lacto-N-fucopentaose II group, a Lacto-N-difucohexaose I group, and a Lactodifucotetraose group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.

29. A synthetic nutrition composition comprising a purified protein modified to include at least two groups selected from: a Lacto-N-fucopentaose I group, a Lacto-N-fucopentaose II group, a 2-Fucosyllactose group, a 3-Fucosyllactose group, a Lacto-N-fucopentaose II group, a Lacto-N-difucohexaose I group, and a Lactodifucotetraose group or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc.

30. A synthetic nutrition composition comprising a purified protein modified to include at least two groups selected from:

Lacto-N-fucopentaose I;

Lacto-N-fucopentaose II;

3'-Fucosyllactose;

Lacto-N-fucopentaose II;

Lacto-N-difucohexaose I;

Lactodifucotetraose;

LactoN-tetraose;

LactoN-neotetraose;

3'-Sialyllactose;

3'-Sialyllactosamine;

20 6'-Sialyllactose;

6'-Sialyllactosamine;

Sialyllacto-N-neotetraose c;

Monosialyllacto-N-hexaose;

Disialyllacto-N-hexaose I;

25 Monosialyllacto-N-neohexaose I;

Monosialyllacto-N-neohexaose II

Disialyllacto-N-neohexaose

Disialyllacto-N-tetraose;

Disialyllacto -N-hexaose II;

30 Sialyllacto-N-tetraose a;

Disialyllacto-N-hexaose I;

Sialyllacto-N-tetraose b;

3'-Sialyl-3-fucosyllactose;

Disialomonofucosyllacto-N-neohexaose;

Monofucosylmonosialyllacto-N-octaose (sialyl Lea);

Sialyllacto-N-fucohexaose II;

Disialyllacto-N-fucopentaose II; and

Monofucosyldisialyllacto-N-tetraose or a variant thereof in which the Glc at the reducing end is replaced with GlcNAc wherein the at least groups are the same or different .

- 31. The composition of claim 30 wherein the protein is modified to include at least two different groups.
  - 32. The synthetic nutritional composition of any of claims 21-29 further comprising an edible fat, a vitamin, a plant protein or an animal protein.
  - 33. A method for treating or reducing the risk of infection, the method comprising administering the composition of any of the forgoing claims wherein said composition is not a mammalian milk.
  - 34. The method of claim 33 wherein the composition comprises 2'FL or 2'FLNAc.
    - 35. The method of claim 34 wherein the molecule comprises a protein to which 2'FL and/or 2'FLNAc are directly or indirectly covalently attached.
  - 36. The method of claim 33 wherein the infection is caused by *V. cholerea* or *C. jejuni*.
    - 37. The method of claim 33 wherein the infection in an enteric infection.

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38. A method for reducing the risk of enteric disease in a patient, the method comprising,

- (a) identifying the two most prevalent agents capable of causing enteric disease in the geographic location of the patient;
- (b) administering to the patient a composition comprising a molecule comprising a first glycan which interferes with the binding to epithelial cells of the first of the two most prevalent agents and a second glycan which interferes with the binding to epithelial cells of the second of the two most prevalent agents wherein said composition is not breast milk.
- 39. A method for reducing the risk of enteric disease in a patient, the method comprising,
  - (a) identifying the two most prevalent agents capable of causing enteric disease in the geographic location of the patient;
    - (b) administering to the patient composition comprising
    - i) a first molecule comprising a first glycan which interferes with the binding to epithelial cells of the first of the two most prevalent agents; and
    - ii) a second molecule glycan which interferes with the binding to epithelial cells of the second of the two most prevalent agents; wherein said composition is not breast milk.
  - 40. A yeast cell harboring a recombinant vector comprising a nucleotide sequence encoding GDP-mannose 4, 6 dehydratase and a nucleotide sequence encoding GDP-L-fucose synthetase.
- 25 41. The yeast cell of claim 40 wherein the GDP-mannose 4, 6 dehydratase is *H. pylori* GDP-mannose 4, 6 dehydratase.
  - 42. The yeast cell of claim 40 or claim 41 wherein the GDP-L-fucose synthetase is *H. pylori* GDP-L-fucose synthetase.

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43. The yeast cell of any of claims 40-42 wherein the yeast cell harbors a nucleic acid molecule encoding a GDP-fucose/GMP antiporter fusion protein.

- 44. The yeast cell of any of claim 43 wherein the fusion protein comprises a golgi-membrane location sequence.
  - The yeast cell of claim 43 wherein the golgi-membrane location sequence is from Vrg4p.
- 46. An isolated nucleic acid molecule encoding a fusion protein comprising at least a first portion and a second portion, the first portion comprising the active domain of a GDP-fucose/GMP antiporter and the second portion comprising a golgi localization sequence.
- 15 47. The isolated nucleic molecule of claim 46 wherein the golgi localization sequence in a yeast golgi localization sequence.
  - 48. A yeast harboring the isolated nucleic acid molecule of claim 46.
- 49. The yeast of claim 48 further harboring a nucleic acid molecule encoding a fucosyltransferase or a galactosyltransferase.
  - 50. The yeast of claim 49 wherein the fucosyltransferase is selected from: Homo sapiens fucosyltransferase 1 (galactoside 2-alpha-L-fucosyltransferase, Bombay phenotype included) (FUT1);

Homo sapiens fucosyltransferase 2 (secretor status included) (FUT2);

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Homo sapiens fucosyltransferase 3 (galactoside 3(4)-L-fucosyltransferase, Lewis blood group included) (FUT3);

Homo sapiens fucosyltransferase 4 (alpha (1,3) fucosyltransferase, myeloid-specific) (FUT4);

Homo sapiens fucosyltransferase 5 (alpha (1,3) fucosyltransferase) (FUT5);

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Homo sapiens fucosyltransferase 6 (alpha (1,3) fucosyltransferase) (FUT6);
Homo sapiens fucosyltransferase 7 (alpha (1,3) fucosyltransferase) (FUT7);
Homo sapiens fucosyltransferase 8 (alpha (1,6) fucosyltransferase) (FUT8);
Homo sapiens fucosyltransferase 9 (alpha (1,3) fucosyltransferase) (FUT9); and
Homo sapiens protein o-fucosyltransferase (POFUT1).
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51. A pharmaceutical composition comprising a purified protein modified to include at least two different groups selected from LNT, LNneoT or a variant thereof wherein the Glc at the reducing end is replaced by GlcNAc.